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(54) Title: PHARMACEUTICAL COMPOSITION COMPRISING AN IMMUNOSUPPRESSANT FOR USE IN THE TREAT-MENT OF SKIN DISEASES

(57) Abstract: Synergistic combinations of a macrolide T-cell immunomodulator or immunosuppressant such as 33-epichloro-33desoxyascomycin and an emollient such as dimethicone, glycerol or isostearyl isostearate are provided, which are useful in particular in the treatment of dermatological or mucosal diseases such as dry skin or atopic or contact dermatitis.



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PHARMACEUTICAL COMPOSITION COMPRISING AN IMMUNOSUPPRESSANT FOR USE IN THE TREATMENT OF SKIN DISEASES

The invention relates to pharmaceutical compositions, for use in particular in the treatment of skin diseases. It concerns a pharmaceutical composition comprising a macrolide T-cell immunomodulator or immunosuppressant and an emollient.

It has now been found that, surprisingly, macrolide T-cell immunomodulators and immunosuppressants, when used in combination or association with emollients, act synergistically, resulting in a potentiation of pharmacological activity, such that effective beneficial, especially anti-dermatitis activity is seen upon co-administration at dosages which would be well below the effective dosages administered individually.

The invention thus concerns novel pharmaceutical compositions comprising a macrolide T-cell immunomodulator or immunosuppressant in association or combination with an emollient, hereinafter briefly named "the compositions of the invention".

A macrolide T-cell immunomodulator or immunosuppressant is to be understood herein as being a T-cell immunomodulator or T-cell immunosuppressant which has a macrocyclic compound structure including a lactone or lactam moiety. While it preferably has at least some T-cell immunomodulating or immunosuppressant activity, it may also exhibit concomitantly or predominantly further pharmaceutical properties, such as anti-inflammatory activity.

An emollient is to be understood herein as being an agent which softens or soothes the skin, or soothes an irritated internal surface.

It should be appreciated that the present invention does not contemplate merely the inclusion of an emollient as a minor excipient in a pharmaceutical composition comprising a macrolide T-cell immunomodulator or immunosuppressant in order to e.g. improve the compatibility of the composition as such with e.g. human skin. More comprehensively, it is contemplated herewith to involve emollients as active agents in their own right, whereby "active" should be understood as relating not only to pharmacological activity, but also activity as regards cosmetic aspects, such as the appearance or brittleness of skin.

The amount of emollient to be used or included with the compositions of the invention is thus normally substantially more than commonly used in pharmaceutical

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compositions, or is administered separately from the macrolide. It is e.g. from about 10 % to about 5000 %, preferably from about 20 % to about 1000 %, more preferably from about 100 % to about 500 % w/w of the amount of macrolide in the composition.

The compositions of the invention may thus be viewed also as health care or personal care products incorporating at least one pharmaceutically active component, or as so-named "cosmeceuticals".

The compositions of the invention may be adapted for systemic use as regards the immunomodulator or immunosuppressant component, e.g. oral or intravenous, or for topical use for both components; preferably they are adapted for topical use. They are useful for the known indications of the particular active agents incorporated therein. They are particularly indicated for use in dermatological or mucosal diseases, e.g. dermatological or mucosal diseases which have an inflammatory component or involve inflammatory complications, such as dry skin or atopic or contact dermatitis.

The composition resulting from the combination is e.g. a medicated emollient, appropriately presented, e.g. as a poultice or a cataplasm.

A suitable macrolide T-cell immunomodulator or immunosuppressant is for example an FKBP12-binding calcineurin inhibitor or mitogen-activated kinase modulator or inhibitor, in particular an asco- or rapamycin. It preferably is an ascomycin. While the macrolide preferably has at least some calcineurin- or mitogen-activated kinase modulating or inhibiting activity, it may also exhibit concomitantly or predominantly further pharmaceutical properties, such as antiinflammatory activity. It preferably is a compound, e.g. an ascomycin, having rather long-acting activity relatively to other members of the same structural class, e.g. it is metabolically degraded slowly to inactive products.

An asco- or rapamycin is to be understood as asco- or rapamycin as such, or a derivative thereof. An asco- or rapamycin derivative is to be understood as being an antagonist, agonist or analogue of the parent compound which retains the basic structure and modulates at least one of the biological, for example immunological properties of the parent compound.

An "anti-inflammatory ascomycin derivative" is defined herein as an ascomycin derivative that exhibits pronounced anti-inflammatory activity in e.g. animal models of allergic

contact dermatitis but has only low potency in suppressing systemic immune response, namely, which has a minimum effective dose (MED) of up to a concentration of about 0.04 % w/v in the murine model of allergic contact dermatitis upon topical administration, while its potency is at least 10 times lower than for tacrolimus (MED 14 mg/kg) in the rat model of allogeneic kidney transplantation upon oral administration (Meingassner, J.G. et al., <u>Br. J. Dermatol.</u> 137 [1997] 568-579; Stuetz, A. <u>Seminars in Cutaneous Medicine and Surgery 20</u> [2001] 233-241). Such compounds are preferably lipophilic.

Suitable ascomycins are e.g. as described in EP 184162, EP 315978, EP 323042, EP 423714, EP 427680, EP 465426, EP 474126, WO 91/13889, WO 91/19495, EP 484936, EP 523088, EP 532089, EP 569337, EP 626385, WO 93/5059 and WO 97/8182; in particular:

- ascomycin;
- tacrolimus (FK506; Prograf^R);
- imidazolylmethoxyascomycin (WO 97/8182 in Example 1 and as compound of formula I);
- 32-O-(1-hydroxyethylindol-5-yl)ascomycin (L-732531) (<u>Transplantation 65</u> [1998] 10-18, 18-26, on page 11, Figure 1; and
- (32-desoxy-32-epi-N1-tetrazolyl)ascomycin (ABT-281) (J.Invest.Dermatol. 12 [1999] 729-738, on page 730, Figure 1); preferably:
- {1R,5Z,9S,12S-[1E-(1R,3R,4R)],13R,14S,17R,18E,21S,23S,24R,25S,27R}-17-ethyl-1,14-dihydroxy-12-[2-(4-hydroxy-3-methoxycyclohexyl)-1-methylvinyl]-23,25-dimethoxy-13,19,21,27-tetramethyl-11,28-dioxa-4-azatricyclo[22.3.1.0(4,9)]octacos-5,18-diene-2,3,10,16-tetraone (Example 8 in EP 626385), hereinafter referred to as "5,6-dehydroascomycin";
- {1E-(1R,3R,4R)]1R,4S,5R,6S,9R,10E,13S,15S,16R,17S,19S,20S}-9-ethyl-6,16,20-trihydroxy-4-[2-(4-hydroxy-3-methoxycyclohexyl)-1-methylvinyl]-15,17-dimethoxy-5,11,13,19-tetramethyl-3-oxa-22-azatricyclo[18.6.1.0(1,22)]heptacos-10-ene-2,8,21,27-tetraone (Examples 6d and 71 in EP 569337), hereinafter referred to as "ASD 732"; and especially

- pimecrolimus (INN recommended) (ASM981; ElidelTM), i.e. {[1E-(1R,3R,4S)]1R,9S,12S, 13R,14S,17R,18E, 21S,23S,24R,25S,27R}-12-[2-(4-chloro-3-methoxycyclohexyl)-1-methylvinyl]-17-ethyl-1,14-dihydroxy-23,25-dimethoxy-13,19,21,27-tetramethyl-11,28,dioxa-4-azatricyclo [22.3.1.0(4,9)]octacos-18-ene-2,3,10,16-tetraone, of formula I

(Example 66a in EP 427680),

hereinafter also referred to as "33-epichloro-33-desoxyascomycin".

Suitable anti-inflammatory ascomycin derivatives are e.g.: (32-desoxy-32-epi-N1-tetrazolyl)ascomycin (ABT-281); 5,6-dehydroascomycin; ASD 732; and pimecrolimus.

Suitable rapamycins are e.g. as described in USP 3'929'992, WO 94/9010 and USP 5'258'389, preferably sirolimus (rapamycin; Rapamune^R) and everolimus (RAD001; Certican^R).

A particularly preferred macrolide T-cell immunomodulator or immunosuppressant is **pimecrolimus**; it is in free form unless specified otherwise.

A suitable emollient is for example one-phase mineral oil (petrolatum), or mineral oil as a two-phase system, either as an oil-in-water or a water-in-oil emulsion, or as a lotion; it is e.g. a silicone such as dimethicone; glycerine; or vaseline. The system may be of low or high viscosity. It may form a hydrophobic protective film on the skin, as with e.g. a silicone such as dimethicone, or paraffin or petrolatum (vaseline). A humectant may be added as appropriate, e.g. glycerol; or an emollient which has semi-occlusive properties may be used,

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such as a fatty acid or a fatty acid ester, e.g. isostearyl isostearate. Preferred emollients are dimethicone, glycerol and isostearyl isostearate.

Emollients may thus be e.g. fatty alcohols, hydrocarbons, triglycerides, waxes, esters, silicone oils and lanolin containing products. Fatty alcohols are e.g. cetyl alcohol, octyldodecanol, stearyl alcohol and oleyl alcohol. Hydrocarbons include mineral oil, petrolatum, paraffin, squalene, polybutene, polyisobuten, hydrogenated polyisobutene, cerisin and polyethylene. Triglycerides are e.g. castor oil, caprylic/capric triglyceride, hydrogenated vegetable oil, sweet almond oil, wheat germ oil, sesame oil, hydrogenated cottonseed oil, coconut oil, wheat germ glycerides, avocado oil, corn oil, trilaurin, hydrogenated castor oil, shea butter, cocoa butter, soybean oil, mink oil, sunflower oil, safflower oil, macadamia nut oil, olive oil, apricot kernel oil, hazelnut oil and borage oil. Waxes include e.g. carnauba wax, beeswax, cadelilla wax paraffin, Japan wax, microcrystalline wax, jojoba oil, cetyl esters wax, and synthetic jojoba oil. Esters include e.g. isopropyl myristate, isopropyl palmitate, octyl palmitate, isopropyl linoleate, C₁₂₋₁₅ alcohol benzoates, cetyl palmitate, myristyl myristate, myristyl lactate, cetyl acetate, propylene glycol dicaprylate/caprate, decyl oleate, stearyl heptanoate, diisostearyl malate, octyl hydroxystearate and isopropyl isostearate. Silicone oils are e.g. dimethicone (dimethyl polysiloxane) and cyclomethicone. Lanolin containing products are e.g. lanolin, lanolin oil, isopropyl lanolate, acetylated lanolin alcohol, acetylated lanolin, hydroxylated lanolin, hydrogenated lanolin and lanolin wax.

Personal care products are e.g. shampoos, hair conditioners, combination shampoo/conditioner, shower gels, soaps, hair styling products, hair colorants, deodorants, antiperspirants and moisturizing lotions. The compositions of the invention may comprise, in addition, further active components which provide benefit to the hair or skin, e.g. moisturizing agents, antiperspirants, anti-bacterials, cleaning agents, hair conditioning agents, hair styling agents, anti-dandruff agents, hair growth promoters, hair dyes and pigments, soaps and perfumes.

The compositions of the invention may be e.g. creamy, of the "light" or "rich" type, or greasy, or containing urea. Further components may be selected from e.g. almond oil, cacao butter, castor oil, decyl oleate, triglyceride, cetostearyl ethylhexanoate, stearyl heptanoate or caprylate, diisopropyl adipate, tri-isononanoin, polyethyleneglycol-40 butyloctanol and trideceth-9, polyethyleneglycol-5-ethylhexanoate.

Subgroups of compositions of the invention comprise a macrolide T-cell immunomodulator or immunosuppressant, preferably an anti-inflammatory ascomycin derivative as defined above, especially pimecrolimus, in combination or association with an emollient other than the following emollients singly or collectively in any number:

- glycerine; and/or
- a fatty acid ester; and/or
- a silicone; and/or
- dimethicone; and/or
- a fatty acid; and/or
- petrolatum.

In a further subgroup of compositions of the invention the macrolide T-cell immunomodulator or immunosuppressant is other than tacrolimus; in a further subgroup it is other than tacrolimus and sirolimus.

Preferred for use in the treatment of conditions where inflammation is involved are compositions of the invention wherein one or both components possess some degree of inherent anti-inflammatory activity. Particularly preferred are compositions comprising an ascomycin, preferably an anti-inflammatory ascomycin derivative, especially pimecrolimus, in combination or association with an emollient; more especially pimecrolimus in combination or association with dimethicone, glycerol or isostearyl isostearate. The inflammatory condition is e.g. dry skin or atopic or contact dermatitis.

Pimecrolimus being anti-inflammatory and having excellent skin penetration but only minimal skin permeation properties, it is not having significant systemic side effects when applied topically on skin, and the soothing effect of emollients complements its anti-inflammatory action on inflamed skin.

"Treatment" as used herein refers in particular to use for preferably alleviating an existing condition, namely curative treatment, although the invention also contemplates prophylactic use in conditions where a high probablity of inflammation exists.

Synergy is e.g. calculated as described in Berenbaum, <u>Clin. Exp. Immunol.</u> 28 (1977) 1, using an interaction term to correct for differences in mechanism between the two

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drugs, as described in Chou et al., <u>Transpl. Proc.</u> 26 (1994) 3043. The index of synergy is calculated as:

$$\frac{\text{dose of A}}{A_E} + \frac{\text{dose of B}}{B_E} + \frac{(\text{dose of A}) \times (\text{dose of B})}{A_E \times B_E}$$

in which the doses of the compounds A and B represent those used in a particular combination, and A_B and B_E are the individual doses of A and B respectively giving the same effect. If the result is less than 1, there is synergy; if the result is 1, the effect is additive; if the result is greater than 1, A and B are antagonistic. By plotting an isobologram of dose of A $/A_E$ vs. dose of B $/B_B$ the combination of maximum synergy can be determined. The synergistic ratio expressed in terms of the ratio by weight of the two compositions at synergistic amounts along the isobologram, especially at or near the point of maximum synergy, can then be used to determine formulations containing an optimally synergistic ratio of the two compounds.

Activity may e.g. be determined in known assay models for testing the activity of the individual components of the compositions.

The invention also provides products and methods for co-administration of a macrolide T-cell immunomodulator or immunosuppressant, e.g. 33-epichloro-33-desoxy-ascomycin or 5,6-dehydroascomycin, and an emollient, e.g. dimethicone, at synergistically effective dosages, e.g.:

- a method of treatment or prevention of a dermatological or mucosal disease such as dry skin or atopic or contact dermatitis in a subject suffering from or at risk for such condition, comprising co-administering synergistically effective amounts of a composition of the invention;
- the use of a macrolide T-cell immunomodulator or immunosuppressant in the manufacture of a medicament for co-administration in synergistically effective amounts with an emollient;
- the use of an emollient in the manufacture of a medicament for co-administration in synergistically effective amounts with a macrolide T-cell immunomodulator or immunosuppressant;
- a kit of parts comprising a macrolide T-cell immunomodulator or immunosuppressant and an emollient in separate unit dosage forms, preferably wherein the unit dosage forms are suitable for administration of the component compounds in synergistically effective amounts, together with instruction for use, optionally with further means for facilitating compliance with the administration of the component compounds, e.g. a label or drawings;

- the use of a macrolide T-cell immunomodulator or immunosuppressant in the manufacture of a pharmaceutical kit which is to be used for facilitating co-administration with an emollient;

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- the use of an emollient in the manufacture of a pharmaceutical kit which is to be used for facilitating co-administration with a macrolide T-cell immunomodulator or immunosuppressant;
- a macrolide T-cell immunomodulator or immunosuppressant and an emollient as a combined pharmaceutical preparation for simultaneous, separate or sequential use, preferably in synergistically effective amounts, e.g. for the treatment or prevention of a dermatological or mucosal disease such as dry skin or atopic or contact dermatitis;
- a pharmaceutical composition comprising a macrolide T-cell immunomodulator or immunosuppressant in combination or association with an emollient, e.g. in synergistically effective amounts, together with at least one pharmaceutically acceptable diluent or carrier, e.g. for use in treatment or prevention of a dermatological or mucosal disease such as dry skin or atopic or contact dermatitis; and
- a process for the preparation of a composition of the invention comprising mixing a macrolide T-cell immunomodulator or immunosuppressant and an emollient, in combination or association with at least one pharmaceutically acceptable diluent or carrier.

By "synergistically effective amounts" is meant an amount of macrolide T-cell immunomodulator or immunosuppressant and an amount of emollient which are individually below their respective effective dosages for a relevant indication, but which are pharmaceutically active on co-administration, e.g. in a synergistic ratio, for example as calculated above. Furthermore, "synergistically effective amounts" may mean an amount of macrolide T-cell immunomodulator or immunosuppressant and an amount of emollient which are individually equal to their respective effective dosages for a relevant indication, and which result in a more than additive effect.

The molar amount of macrolide T-cell immunomodulator or immunosuppressant present is from roughly similar to, to significantly less than the amount of emollient, preferably half as much or less. Synergistic ratios of macrolide T-cell immunomodulator or immunosuppressant to emollient by weight are thus suitably from about 10:1 to about 1:50, preferably from about 5:1 to about 1:20, most preferably from about 1:1 to about 1:15, e.g. about 1:12.

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The compositions of the invention can be administered as a free combination, or can be formulated into a fixed combination, which greatly enhances the convenience for the patient.

Absolute dosages of the compounds will vary depending on a number of factors, e.g. the individual, the route of administration, the desired duration, the rate of release of the active agent and the nature and severity of the condition to be treated. For example, the amount of active agents required and the release rate thereof may be determined on the basis of known in vitro and in vivo techniques, determining how long a particular active agent concentration in the blood plasma remains at an acceptable level for a therapeutic effect.

For example, in prevention and treatment of a dermatological or mucosal disease such as dry skin or atopic or contact dermatitis, an initial dosage of about 2-3 times the maintenance dosage is suitably administered, followed by a daily dosage of about 2-3 times the maintenance dosage for a period of from one to two weeks, and subsequently the dose is gradually tapered down at a rate of about 5 % per week to reach the maintenance dosage.. In general, synergistically effective amounts of 33-epichloro-33-desoxyascomycin and dimethicone on oral administration for use in prevention and treatment of dry skin or atopic or contact dermatitis in larger animals, e.g. man, are amounts of pimecrolimus of up to about 2 mg/kg/day, e.g. from about 0.01 mg/kg/day to about 2 mg/kg/day, preferably about 0.5 mg/kg/day, in combination or co-administration with amounts of dimethicone of up to about 50 mg/kg/day, e.g. from about 0.25 mg/kg/day to about 50 mg/kg/day, preferably about 2.5 mg/kg/day, in a synergistic ratio, as described. Suitable unit dosage forms for oral co-administration of these compounds thus may contain on the order of from about 0.5 mg to about 100 mg, preferably about 3 mg to about 30 mg of 33-epichloro-33-desoxyascomycin, and from about 10 mg to about 3000 mg, preferably about 50 mg to about 500 mg of dimethicone. The daily dosage for oral administration is preferably taken in a single dose, but may be spread out over two, three or four dosages per day. For i.v. administration, the effective dosage is lower than that required for oral administration, e.g. about one fifth the oral dosage.

By "co-administration" is meant administration of the components of the compositions of the invention together or at substantially the same time, e.g. within fifteen minutes or less, either in the same vehicle or in separate vehicles, so that upon oral

administration, for example, both compounds are present simultaneously in the gastrointestinal tract. However, upon topical application, administration of the components may also be separated by a time interval of at least several hours, e.g. 6 hours or 12 hours. Preferably, the compounds are administered as a fixed combination, preferably topically.

The compositions of the invention include compositions suitable for administration by any conventional route, in particular compositions suitable for administration either enterally, for example, orally, e.g. in the form of solutions for drinking, tablets or capsules, or parenterally, e.g. in the form of injectable solutions or suspensions; or topically, e.g. for the treatment of inflammatory conditions of the skin or mucosae, e.g. in the form of a dermal cream, ointment, ear drops, mousse, shampoo, solution, lotion, gel, emulgel or like preparation, e.g. in a concentration of from about 0.1 % to about 2 %, preferably about 1 % by weight of each component, especially in combination or association with penetration enhancing agents, as well as for application to the eye, e.g. in the form of an ocular cream, gel or eye-drop preparation, for treatment of inflammatory conditions of the lungs and airways, e.g. in the form of inhalable compositions, and for mucosal application, e.g. in the form of vaginal tablets.

The compositions of the invention are suitably emulsions, microemulsions, emulsion preconcentrates or microemulsion preconcentrates, or solid dispersions, especially water-in-oil microemulsion preconcentrates or oil-in-water microemulsions, comprising the macrolide T-cell immunomodulator or immunosuppressant and the emollient in a synergistic ratio.

The compositions of the invention can be prepared in conventional manner, e.g. by mixing a macrolide T-cell immunomodulator or immunosuppressant and an emollient, in combination or association with at least one pharmaceutically acceptable diluent or carrier.

The active agent components may be in free form or pharmaceutically acceptable salt form as appropriate.

While the invention primarily contemplates combination or association of just two pharmaceutically and/or cosmetically active components, it does not exclude the presence of further pharmaceutically and/or cosmetically active agents, e.g. one further active agent, such as an antiseptic, as far as they do not contradict the purpose of the present invention.

The following Examples illustrate the invention. The compounds are in free, i.e. neutral or base form unless specified otherwise.

Example 1: Cream (protective hydrophobic film)

Component	Amount (g)	•
33-Epichloro-33-desoxyascomycin	1.00	
dimethicone	5.00	
triglycerides, medium chain	15.00	
oleyl alcohol	10.00	
sodium cetylstearyl sulfate	1.00	
cetyl alcohol	4.00	
stearyl alcohol	4.00	
glyceryl monostearate	2.00	
penzyl alcohol	1.00	
propylene glycol	5.00	•
citric acid	0.05	
sodium hydroxide	*	
•	d 100.0	
amount required to adjust pH to 5	.5	

Preparation is according to conventional manufacturing procedures for an emulsion. The ascomycin derivative and dimethicone are added to the heated homogeneous oily phase which contains triglycerides medium chain, oleyl alcohol, sodium cetylstearyl sulfate, cetyl alcohol, stearyl alcohol and glyceryl monostearate. In parallel, the water phase containing the remaining ingredients is heated at the same temperature as the oily phase. The oily phase is added to the water phase and homogeneisation is performed. The resultant cream is cooled to room temperature.

Example 2: Cream (with a humectant)

The composition is as for Example 1, whereby dimethicone 5.00 g is replaced with glycerol 3.00 g, which for preparation is included in the water phase in place of the oily phase.

Example 3: Cream (semi-occlusive)

The composition is as for Example 1, whereby dimethicone 5.00 g is replaced with isostearyl isostearate 4.00 g.

Example 4: Ointment (protective hydrophobic film)

Component	Amount (g)	
33-Epichloro-33-desoxyascomycin	1.00	
dimethicone	5.00	
oleyl alcohol	10.00	
hexylene glycol	10.00	
microcrystalline wax	5.00	
white vaseline	ad 100.0	

Preparation is according to conventional manufacturing procedures. The ascomycin is added to the heated homogeneous oily phase which contains dimethicone and the remaining ingredients. After homogeneisation the resultant ointment is cooled to room temperature.

Example 5: Solution (protective hydrophobic film)

Component	Amount (g)	
33-Epichloro-33-desoxyascomycin	1.00	
dimethicone	5.00	
triglycerides, medium chain	10.00	
oleyl alcohol	10.00	
	d 100.0	

Preparation is according to conventional manufacturing procedures and is as described under Example 4.

Example 6: Liquid emulsion (with a humectant)

Component	Amount (g)	•
33-Epichloro-33-desoxyascomycin	1.00	
glycerol	3.00	•••
triglycerides, medium chain	15.00	
oleyl alcohol	10.00	
glyceryl monooleate	2.00	
Tween 80	4.00	
benzyl alcohol	1.00	
propylene glycol	5.00	
citric acid	0.05	
sodium hydroxide	*	
	d 100.0	

Preparation is according to conventional manufacturing procedures. The ascomycin is added to the heated homogeneous oily phase which contains triglycerides medium chain, oleyl alcohol and glyceryl monooleate. In parallel, the water phase containing glycerol and the remaining ingredients is heated at the same temperature as the oily phase. The oily phase is added to the water phase and homogeneisation is performed. The resultant emulsion is cooled to room temperature.

Example 7: Liquid emulsion (semi-occlusive)

As for Example 6, whereby glycerol 3.00 g is replaced with isostearyl isostearate 4.00 g, which for preparation is included in the oily phase in place of the water phase.

Claims:

- 1. A pharmaceutical composition comprising a macrolide T-cell immunomodulator or immunosuppressant in combination or association with an emollient, together with at least one pharmaceutically acceptable diluent or carrier.
- 2. A composition according to claim 1 comprising 33-epichloro-33-desoxyascomycin in combination or association with dimethicone, glycerol or isostearyl isostearate.
- 3. A method of treatment of a dermatological or mucosal disease such as dry skin or atopic or contact dermatitis in a subject suffering from or at risk for such condition, comprising co-administering a synergistically effective amount of a composition according to claim 1.
- 4. A process for the preparation of a composition according to claim 1 comprising mixing a macrolide T-cell immunomodulator or immunosuppressant and an emollient, in combination or association with at least one pharmaceutically acceptable diluent or carrier.
- 5. A kit of parts comprising a macrolide T-cell immunomodulator or immunosuppressant and an emollient in separate unit dosage forms, together with instructions for use.

INTERESTIONAL SEARCH REPORT

International Application No
PCT/EP2004/003513

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 A61K31/435 A61K31/436 A61K47/02 A61K47/10 A61K47/14 A61K47/44 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) A61K A61P IPC 7 Documentation searched other than minimum documentation to the extent that such documents are included. In the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, BIOSIS, EMBASE, WPI Data, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category * Relevant to claim No. WO 96/13249 A (SANDOZ AG; SCHMOOK FRITZ 1-5 (AT); POPP XUE PING (CH); JACKMAN MARTIN (CH) 9 May 1996 (1996-05-09) page 1 - page 15 WO 00/32234 A (NOVARTIS ERFIND VERWALT X 1-5 GMBH; NOVARTIS AG (CH); KRIWET KATRIN (DE); R) 8 June 2000 (2000-06-08) page 1 - page 13 Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the International filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention *E* earlier document but published on or after the International "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to *L* document which may throw doubts on priority claim(s) or involve an inventive step when the document is taken alone which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the *O* document referring to an oral disclosure, use, exhibition or document is combined with one or more other such documents, such combination being obvious to a person skilled other means In the art. *P* document published prior to the International filing date but later than the priority date claimed *&* document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 25/08/2004 18 August 2004 Name and mailing address of the ISA Authorized officer European Palent Office, P.B. 5818 Palentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nt, Kling, I Fax: (+31-70) 340-3016

INTERESTIONAL SEARCH REPORT

	CROND OCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	KAPP A ET AL: "LONG-TERM MANAGEMENT OF ATOPIC DERMATITIS IN INFANTS WITH TOPICAL PIMECROLIMUS, A NONSTEROID ANTI-INFLAMMATORY DRUG" JOURNAL OF ALLERGY AND CLINICAL IMMUNOLOGY, MOSBY - YEARLY BOOK, INC, US, vol. 110, no. 2, August 2002 (2002-08), pages 277-284, XP009032310 ISSN: 0091-6749 the whole document	1-5
X	WO 97/25977 A (CIBA GEIGY AG; TIEMESSEN HARRY (DE)) 24 July 1997 (1997-07-24) page 1 - page 16	1-5
X	EP 0 812 588 A (YOSHITOMI PHARMACEUTICAL) 17 December 1997 (1997-12-17) the whole document	1-5
X	EP 1 273 288 A (NOVARTIS ERFIND VERWALT GMBH; NOVARTIS AG (CH)) 8 January 2003 (2003-01-08) page 2, paragraph 1 - page 7, paragraph 3	1-5
X	GB 2 327 610 A (NOVARTIS AG) 3 February 1999 (1999-02-03) page 1 - page 5	1-5
Υ .	EP 1 064 942 A (FUJISAWA PHARMACEUTICAL CO) 3 January 2001 (2001-01-03) page 2 - page 11	1-5
Y	US 2002/044967 A1 (IBUKI RINTA ET AL) 18 April 2002 (2002-04-18) page 1, left-hand column - page 8, left-hand column	1-5
P,X	WO 2004/016289 A (NOVARTIS PHARMA GMBH; NOVARTIS AG (CH); SEKKAT NABILA (CH); KRIWET KA) 26 February 2004 (2004-02-26) the whole document	1-5
P,X	WO 03/074054 A (NOVARTIS PHARMA GMBH; NOVARTIS AG (CH); BABIOLE SAUNIER MAGGY (FR); B) 12 September 2003 (2003-09-12) the whole document	1-5

information on patent family members

	atent document I In search report		Publication date		Patent family member(s)		Publication date
WO	9613249	Α	09-05-1996	AT	214593	T	15-04-2002
				AU	714254		23-12-1999
				AU	3845195		23-05-1996
				AU	5833699	Α	06-01-2000
				BR	9509530	Α	14-10-1997
				CA	2200966	A1	09-05-1996
				CN	1401325	=	12-03-2003
				CN	1162259		15-10-1997
				CY	2211		08-11-2002
				CZ	9701232	A3	13-08-1997
	•			·CZ		B6	12-06-2002
				DE	19581804		22-01-1998
		•		DE		D1	25-04-2002
				DE	69525957		14-11-2002
				DK	786986		29-04-2002
	•			MO	9613249		09-05-1996
				EP	1147766		24-10-2001
				EP	0786986		06-08-1997
	•			ES	2173978		01-11-2002
	•			FI	971018		18-04-1997
				GB	2308546		02-07-1997
				HU	77140		02-03-1998
				JP NO	10508588		25-08-1998
				NZ	971951 <i>(</i> 295170 <i>(</i>	-	25-04-1997 25-02-1000
				NZ	331824		25-02-1999
			•	PL	331824 <i>1</i> 319599 <i>1</i>		28-01-2000
			•	PL	185320 I		18-08-1997 30-04-2003
				PT	786986		31-07-2002
				SI	786986 .1	•	31-07-2002
				SK	52097 <i>i</i>		10-09-1997
	•			US	2001031769	· -	18-10-2001
			·	CY	2212		08-11-2002
•				GB	2327610		03-02-1999
				PL	184908	•	31-01-2003
نغ، سنة بالشد	ه من بعرده الله کامل بسمه به سات	يونون سن جب جب تب	ن کا کا مناسا من کری چرد ساخا سا کا کا د	RU	2181592	C2	27-04-2002
WO	0032234	Α	08-06-2000	AU	767156		30-10-2003
				AU	1656900	•	19-06-2000
				BR	9915861		21-08-2001
				CA	2350884		08-06-2000
				CN	1329507	_	02-01-2002
				CZ	20011908	· -	12-09-2001
				WO	0032234 /		08-06-2000
				EP	1135163		26-09-2001
				HU	0104413 /	42 r	28-03-2002
				JP NO	2002531419	l A	24-09-2002
				NO NZ	20012624 /		09-07-2001
				NZ Pl	511687 /		31-10-2003
				SK	348750 / 7622001 /		03-06-2002
				TR	200101547		06-11-2001
				US	200101547		22-10-2001 13-12-2001
				ZA	2001031030 /	· -	04-06-2002
WO	9725977	 А	24-07-1997	 AT	220440	 r	هن ها، هروه هنا الآيا هنا هي جند هنا هنا هجا گذا ك
		П	LT U/-133/	AU	239449] 1543497 <i> </i>	=	15-05-2003
				CA	2240339 <i>f</i>		11-08-1997 24-07-1997
				JA	EE-10303 1	1.7	74-0/-133/

Information on patent family members

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 9725977	A		DE DE DK WO EP EP ES HK JP PT US	69721729 D1 69721729 T2 874621 T3 9725977 A1 1273288 A1 1273289 A1 0874621 A1 2199338 T3 1015277 A1 2000503655 T 2004107350 A 874621 T 6239102 B1	12-06-2003 11-12-2003 01-09-2003 24-07-1997 08-01-2003 08-01-2003 04-11-1998 16-02-2004 08-04-2004 28-03-2000 08-04-2004 30-09-2003 29-05-2001
EP 0812588	A	17-12-1997	AU AU EP IL NZ SK US CZ HU WO RU US	705320 B2 1172997 A 0812588 A1 121625 A 324453 A 115797 A3 6121329 A 2213989 A1 9702704 A3 9800034 A2 9724112 A1 2156127 C2 6197829 B1	20-05-1999 28-07-1997 17-12-1997 13-09-2001 27-03-2000 04-02-1998 19-09-2000 10-07-1997 14-01-1998 28-05-1999 10-07-1997 20-09-2000 06-03-2001
EP 1273288	A	08-01-2003	EP EP AU CDE DK OP ES HP JP US	1273288 A1 1273289 A1 239449 T 1543497 A 2240339 A1 69721729 D1 69721729 T2 874621 T3 9725977 A1 0874621 A1 2199338 T3 1015277 A1 2000503655 T 2004107350 A 874621 T 6239102 B1	08-01-2003 08-01-2003 15-05-2003 11-08-1997 24-07-1997 12-06-2003 11-12-2003 01-09-2003 24-07-1997 04-11-1998 16-02-2004 08-04-2004 28-03-2000 08-04-2004 30-09-2003 29-05-2001
GB 2327610	A	03-02-1999	GB AU AU AU BR CN CN CY CY CZ DE	2308546 A ,B 214593 T 714254 B2 3845195 A 5833699 A 9509530 A 2200966 A1 1401325 A 1162259 A ,B 2211 A 2212 A 9701232 A3 19581804 T0	02-07-1997 15-04-2002 23-12-1999 23-05-1996 06-01-2000 14-10-1997 09-05-1996 12-03-2003 15-10-1997 08-11-2002 08-11-2002 13-08-1997 22-01-1998

INTERATIONAL SEARCH REPORT

Information on patent family members

·	,			1.01	7 21 2004/ 000013
Patent document cited in search report		Publication date		Patent family member(s)	Publication date
. GB 2327610 A	Δ		DE	69525957 D1	25-04-2002
			DE	69525957 T2	. — – –
			DK	786986 T3	
			WO	9613249 A1	09-05-1996
			EP	1147766 A2	
			EP	0786986 A1	_ ·
			ES	2173978 T3	
			FI	-	
				971018 A	18-04-1997
			UH	77140 A2	
			JP	10508588 T	25-08-1998
			NO	971951 A	25-04-1997
			NZ	295170 A	25-02-1999
			NZ	331824 A	28-01-2000
			PL	319599 A1	18-08-1997
			PL	184908 B1	
			PT	786986 T	31-07-2002
	•		RU	2181592 C2	
			SI	786986 T1	30-06-2002
			SK	52097 A3	
ے بہر سے سے جے جے بنیاضا انگا انتخاب کے سا		···· —	US 	2001031769 A1	18-10-2001
EP 1064942	Α	03-01-2001	AT	269075 T	15-07-2004
			AU	749623 B2	27-06-2002
			AU	2856399 A	18-10-1999
			BR	9909201 A	14-11-2000
			CA	2322516 A1	07-10-1999
			DE	69918074 D1	22-07-2004
			EP	1064942 A1	03-01-2001
			HR	20000707 A1	
			HU	0101237 A2	
•			NO	20004773 A	23-11-2000
			NZ	507211 A	25-07-2003
		•	RU	2214244 C2	
			SK	14392000 A3	
		•	US	6440458 B1	27-08-2002
			CN	1301157 T	27-06-2001
			EP	1421939 A1	26-05-2004
•			ID	27825 A	26-04-2001
			WO	9949863 A1	07-10-1999
			PL	343096 A1	30-07-2001
			TR	200002771 T2	. 21-02-2001
			TW	570814 B	11-01-2004
			US	2003235614 A1	25-12-2003
			US	2003233014 A1 2002044967 A1	18-04-2002
			ZA	2002044967 AT 200004963 A	
	·		<i></i>		08-01-2002
US 2002044967	A1	18-04-2002	US	2003235614 A1	25-12-2003
		•	AT	269075 T	15-07-2004
			AU	749623 B2	27-06-2002
•			AU	2856399 A	18-10-1999
			BR	9909201 A	14-11-2000
			CA	2322516 A1	07-10-1999
			CN	1301157 T	27-06-2001
			DE	69918074 D1	22-07-2004
					* *
			EP	1421939 A1	26-05-2004
			EP	1064942 A1	26-05-2004 03-01-2001
				· · · · · · ·	



Information on patent family members

Patent document cited in search report	Publication date		Patent family member(s)		Publication date
US 2002044967	A1	ID	27825	A	26-04-2001
		WO	9949863	A1	07-10-1999
		NO	20004773	Α	23-11-2000
		NZ	507211	Α	25-07-2003
		PL	343096	A1	30-07-2001
		SK	14392000	A3	12-03-2001
		TR	200002771	T2	21-02-2001
		TW	570814	В	11-01-2004
		US	6440458	B1	27-08-2002
		ZA	200004963	Α	08-01-2002
WO 2004016289	A 26-02-2004	WO	2004016289	A1	26-02-2004
WO 03074054	A 12-09-2003	WO	03074054	A1	12-09-2003

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- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
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- with amended claims

Date of publication of the amended claims: 6 January 2005

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: PHARMACEUTICAL COMPOSITION COMPRISING AN IMMUNOSUPPRESSANT FOR USE IN THE TREATMENT OF SKIN DISEASES

(57) Abstract: Synergistic combinations of a macrolide T-cell immunomodulator or immunosuppressant such as 33-epichloro-33-desoxyascomycin and an emollient such as dimethicone, glycerol or isostearyl isostearate are provided, which are useful in particular in the treatment of dermatological or mucosal diseases such as dry skin or atopic or contact dermatitis.



2004/087141 A1

AMENDED CLAIMS

[received by the International Bureau on 15 October 2004 (15.10.04); original claims 1, 2, 4, 5 replaced by amended claims 1, 2, 4, 5]

Patent Claims

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- 1. A pharmaceutical composition comprising 33-epichloro-33-desoxyascomycin in combination or association with an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate together with at least one pharmaceutically acceptable diluent or carrier.
- 2. A pharmaceutical composition of claim 1 wherein the emollient is present in an amount from about 10% to about 5000% w/w of the amount of 33-epichloro-33-desoxyascomycin.
- A method of treatment of a dermatological or mucosal disease in a subject suffering from such a disease comprising co-administering synergistically effective amounts of a composition of claim
 1.
- 4. A process for the preparation of a composition of any one of claims 1 or 2 comprising mixing 33-epichloro-33-desoxyascomycin and an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate in combination or association with at least one pharmaceutically acceptable diluent or carrier.
- 5. A kit of parts comprising 33-epichloro-33-desoxyascomycin and an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate in separate unit dosage forms, together with instructions for use.